## **PCT**

# WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



#### INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:

C11D 17/00, A61K 7/06, 7/48, C11D 3/00

(11) International Publication Number:

WO 98/28398

A1 |

(43) International Publication Date:

2 July 1998 (02.07.98)

(21) International Application Number:

PCT/GB97/03530

(22) International Filing Date:

23 December 1997 (23.12.97)

(30) Priority Data:

96309466.9

23 December 1996 (23.12.96) EP

(34) Countries for which the regional or

international application was filed: GB et al.

(71) Applicants: QUEST INTERNATIONAL B.V. [NL/NL]; Huizerstraatweg 28, NL-1411 GP Naarden (NL). ALLIED COLLOIDS LIMITED [GB/GB]; Cleckheaton Road, Low Moor, P.O. Box 38, Bradford, West Yorkshire BD12 0JZ (GB).

(72) Inventors: NESS, Jeremy, Nicholas; 25 High Snoad Wood, Challock, Ashford, Kent TN25 4DQ (GB). IRVING, Pamela, Virginia; 8 Calgary Crescent, Folkestone, Kent CT19 6JA (GB). GOODALL, Marcus, James; 102 Cradlebridge Drive, Willesborough, Ashford, Kent TN24 0RF (GB). SYMES, Kenneth, Charles; 4 Silk Mill Drive, East Moreton, Keighley, Yorkshire BD20 5UU (GB). GREY, Bryan, David; 48 Cleckheaton Road, Bradford, West Yorkshire BD6 1BE (GB).

(74) Agents: HARRISON, David, C. et al.; Mewburn Ellis, York House, 23 Kingsway, London WC2B 6HP (GB). (81) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, GM, GW, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG).

#### **Published**

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: PARTICLES CONTAINING ABSORBED LIQUIDS AND METHODS OF MAKING THEM

#### (57) Abstract

Perfume is absorbed within organic polymer particles which have a further polymer at their exterior. The further polymer incorporates free hydroxyl groups and serves to promote deposition of the particles from a wash or rinse liquor. The further polymer may be part of an encapsulating shell, but more conveniently is used as a stabiliser during polymerisation of the particles. Highly hydrolysed polyvinyl alcohol is preferred.

# FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
AU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
AZ	Azerbaijan	GB	United Kingdom	MC	Мопасо	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	т	Trinidad and Tobago
BJ	Benin	IE	Ireland	MN	Mongolia	ÜA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of Americ
CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NB	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	ZW	Yugoslavia Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand	2.44	Zimoaowe
CM	Cameroon	•	Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korca	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	LI	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		

1

# PARTICLES CONTAINING ABSORBED LIQUIDS AND METHODS OF MAKING THEM

This invention relates to particles containing absorbed liquids. An example of such a liquid is a perfume; another is a sunscreening agent. They are useful in for example detergent, personal care, cosmetic and sunscreening products.

10

25

#### Summary of the Prior Art

There have been a number of proposals for absorbing perfume onto a solid carrier material, and various reasons for doing so. In a number of instances, the objective of such proposals has been merely to convert liquid perfume into a solid form which can more readily be incorporated into another product. It has been asserted that such carrier substances may serve to protect the perfume from loss during storage or protect it from contact with other constituents of a composition.

Examples of disclosures of such carriers for perfume are GB-A-1306924, US-A-3449266, US-A-3909461, US-A-4536315, US-A-4539135, US-A-4713193, GB-A-2066839, EP-A-332259, EP-A-332260 and JP-A-63/79662. In many of these documents the carriers are inorganic materials.

EP-A-535942 and EP-A-536942 are concerned with inorganic carrier materials which serve to reduce the vapour pressure of absorbed perfume.

2

As pointed out in those documents, a carrier material which reduces the perfume vapour pressure over the solid carrier but releases the perfume on contact with water can be useful to avoid giving excessive perfume odour to a concentrated product, yet provide the required concentration of perfume in a wash or rinse liquor.

EP-A-285694 is one of a series of documents concerned with porous cross linked polymers obtained by polymerisation around droplets of a liquid substance which may subsequently be removed. These porous polymers can act as carriers for a variety of liquids. They are used in products for application to the skin, and give controlled release of the liquid to the skin.

US 4842761 discloses composite particles in which one polymer is embedded within another. These particles are used as perfume carriers in laundry detergents and provide controlled release of perfume.

#### 20 The invention

We have now found that the retention or controlled release of liquids from particles can be improved by imbibing the liquid into particles containing organic polymer having at the surface of the particles a further polymer with free hydroxyl groups. Such polymer enhances deposition of the particles and by doing so also enhances deposition of the imbibed liquid. The use of such particles can enhance deposition of the liquid, it can

3

retard the evaporation of deposited liquid and can also enhance the extent to which deposited liquid survives a subsequent drying step.

Accordingly, the present invention provides

5 particles containing organic polymer, which are insoluble in water, with liquid imbibed by the particles, the particles having at their exterior, a polymer which incorporates free hydroxy groups. This polymer will be referred to below as a "hydroxy functional polymer".

The invention also provides products which are compositions or articles containing particles according to the invention together with other ingredients. The quantity of the particles will depend on the nature of the product but will generally provide liquid in an amount within the broad range 0.1 to 10% by weight. In particular such products may be personal washing compositions, fabric detergent compositions, other cleaning compositions, fabric conditioner compositions, "leave-on" hair and skin care compositions or sunscreening compositions incorporating such liquid-

As will be explained in greater detail below, the organic polymer particles are solid (but possibly porous solid) particles into which liquid is absorbed by imbibition. Polymer capsules may enclose a solid core into which perfume is absorbed.

containing particles.

25

The particles themselves are insoluble in water, so that they can be deposited from an aqueous wash or rinse

4

liquor.

The hydroxy functional polymer at the exterior of these particles may form, or be included within a coating or incomplete coating on these individual particles. The hydroxy functional polymer is preferably nonionic or cationic. It will be explained below that it generally constitutes between 1 and 25% of the weight of the particles, usually between 1 and 10%.

Cationic functionalities may additionally be present

10 at the exterior of the particles, and may be provided by
the hydroxy functional polymer or otherwise. Presence of
cationic monomers has a tendency to increase particle
size.

#### 15 Particle size

20

Polymer particles used in this invention desirably have an average particle size of at least  $10\mu\text{m}$ , better at least  $20\mu\text{m}$  or  $30\mu\text{m}$ , for ease of handling. Also, we have observed that the rate of release of liquid may be faster than desired if the particles are of very small size such as average size of  $1\mu\text{m}$ . The polymer particles desirably have an average size not larger than  $150\mu\text{m}$ , better not over  $125\mu\text{m}$  so that the particles are not easily visible after deposition.

For particles intended to be used in fabric washing or conditioning, it is especially preferred to use particles with a mean size of at least  $40\,\mu\text{m}$ , e.g. 40 to  $100\,\mu\text{m}$  to retain perfume in the particles and provide slow

5

liquid release.

For particles intended for other products, e.g. personal washing products, a faster rate of release may be desired than with fabrics products, although retention of liquid and delay in release for some hours is still desirable, consequently a smaller particle size may be advantageous, such as a mean size in the range from 10 to  $50\mu m$ .

Polymerisation techniques generally produce a range of particle sizes. For this invention it is desirable that a high proportion of the particles lie between the above limits on particle size. Thus, when particles are intended for fabric treatment, preferably 90% or more of the particles are larger than 30μm. Preferably also 90% or more of the particles are not larger than 150 or even 125μ. Better, 95% or more are not larger than 125 or even 100μm.

To achieve these criteria, it may be necessary to sieve the particles and thereby separate oversized and/or undersized particles. An important aspect of reducing the amount of emulsion polymer formed (broadly speaking, polymer beads of a size below 10 \mum) is to use an initiator system that is substantially insoluble in the aqueous phase, so as to prevent initiation of monomer molecules held in micelles. Suitable water insoluble initiators include azo compounds such as azobisisobutyronitrile (AIBN) and higher alkyl peroxides.

### Hydroxy functional polymer

20

This polymer bearing hydroxy groups and located at the exterior of the particles serves to enhance deposition onto (or retention on) skin, hair, hard surfaces especially vitreous surfaces and fabric.

This polymer is desirably such that at least 80% of the monomer residues in the polymer contain at least one free hydroxy group able to participate in hydrogen bonding. The polymer is preferably nonionic and such that, in its pure state, it is water-soluble. However, when attached to the exterior of the particles, it does not render them water soluble. The attachment is such that the polymer is not completely removed upon contact with water. It may be chemically bonded with the core polymer.

The preferred polymer is polyvinyl alcohol.

As is well known, polyvinyl alcohol cannot be prepared directly by polymerisation, and is obtained by the hydrolysis of polyvinyl acetate. This hydrolysis generally stops before completion, and polymers with varying amounts of hydrolysis are commercially available.

We have found that deposition of particles, especially onto fabric, is better when the polyvinyl alcohol exhibits a substantial level of hydrolysis.

We may therefore prefer to use polyvinyl alcohol which is 85 to 100% hydrolysed i.e. in which 85 or 88 to 100% of the acetate residues have been removed by hydrolysis. More preferably, the polyvinyl alcohol is at

7

least 90% better at least 96% hydrolysed, e.g. 96 to 99% hydrolysed.

Another possible polymer is a copolymer of polyvinyl alcohol, available through hydrolysis of the

5 corresponding copolymer of vinyl acetate. For example a copolymer of vinyl acetate and vinyl formamide, with a high proportion of vinyl acetate can by hydrolysed to a copolymer of vinyl alcohol and vinyl formamide. Further hydrolysis of such a polymer converts the formamide

10 residues to amine groups, giving a cationic copolymer of vinyl alcohol and vinyl amine.

Another category of polymers which can be used is cellulose and chemically modified cellulose where the modification does not greatly reduce the number of hydroxyl groups present. Examples of polymers in this category are hydroxyethyl cellulose, and hydroxypropyl cellulose.

Hydroxyethyl cellulose is available commercially and is made by treatment of cellulose with sodium hydroxide,

20 followed by reaction with ethylene oxide. Another possibility is cellulose which has been chemically modified by the introduction of cationic substituent groups.

It is a characteristic of many grades of polyvinyl alcohol and of hydroxy ethyl cellulose that free hydroxy groups constitute more than 20%, often more than 25% of the weight of the polymer.

#### Solid particles

This invention uses polymer particles which are solid - although they may be porous as well as solid - rather than particles in the form of hollow capsules.

Advantages of solid particles over capsules are that the desired size range is accessible, and that the polymerisation reaction can be carried out in the absence of the liquid.

The particles could possibly be porous particles

10 made by polymerisation around a liquid pore-forming

agent, as taught in EP-A-285694. However, we have found

it to be unnecessary to include such a pore-forming

agent. Polymers formed by simple polymerisation of a

monomer mixture are able to absorb and carry perfume.

15 Preferred polymer particles may be formed by the polymerisation of vinyl monomers, with some cross linking and/or chain branching agent included in the monomers which are polymerised, so that some cross links are formed between the polymer chains. If a cross linking 20 agent is used, the proportion of cross linking may be low, so that after polymerisation there may be some polymer chains which remain entirely linear and are not cross linked to any other chains.

A number of vinyl monomers containing a single

25 carbon-carbon double bond may be used. One suitable
category of monomers is the esters of acrylic and alkyl
acrylic acids, of formula

9

 $H_2C = C$   $CO_2R_2$ 

where  $R_1$  is hydrogen or straight or branched alkyl of 1 to 6 carbon atoms, preferably 1 to 3 carbon atoms and  $R_2$  is straight or branched alkyl of 1 to 6 carbon atoms, preferably 3 to 6 and most preferably 3 or 4 carbon atoms in a straight or branched chain.

Specific examples of suitable monomers are isobutyl

methacrylate (which is particularly preferred), n-butyl
acrylate, n-butyl methacrylate, isobutyl acrylate, npropyl acrylate and iso-propylmethacrylate. Less
preferred is methyl methacrylate.

Another suitable monomer is styrene.

It is possible to use simple linear polymers.

However, these can give particles which are somewhat sticky, and it is usually convenient to introduce some cross-linking or chain branching.

above monomers can be achieved by including in the monomer mixture a small proportion - for example less than 10%, preferably as little as 5% or 1% - of a monomer with two carbon-carbon double bonds. The use of such a material to provide cross linking is well known in other applications of polymers, although it is usual to introduce a greater proportion of cross linking than is required for this invention. Examples of this type of cross linking agent are divinyl benzene, diesters formed between acrylic acid and diols, such as 1,4 butane diol

diacrylate, and higher esters formed between acrylic acid and polyols - which may be sugars.

Chain branching can be introduced by including among the monomers a hydroxyalkyl monomer of formula

5

25

11

 $H_2C = C$   $CO_2R_2$ 

where R<sub>1</sub> is as specified above and R<sub>3</sub> is alkyl of 1 to 6 carbon atoms bearing at least one hydroxy group, preferably 3 to 4 carbon atoms in a straight or branched 10 chain and bearing a single hydroxy group. These monomers undergo a side reaction during the course of polymerisation, and this side reaction produces chain branching. When there is chain branching without cross linking, it is suitable that hydroxyalkyl monomer of the above formula provides from 10 to 40% by weight of the monomer mixture.

Suitable hydroxyalkyl acrylate monomers are hydroxypropyl methacrylate, hydroxybutylacrylate, and hydroxyethylacrylate.

Attachment of a polymer with hydroxy groups, notably polyvinyl alcohol, at the exterior of the particles, can be achieved by polymerising the monomers in the presence of the polyvinyl alcohol (or other polymer with hydroxy groups) using the technique of suspension polymerisation.

Suspension polymerisation is a process in which the organic monomers are formed into a suspension in an aqueous phase, and polymerised. Polymerisation of each suspended droplet leads to a bead of polymer.

It is customary to stabilise the suspension by incorporating a stabilising agent in the aqueous phase

before adding the monomer. Polyvinyl alcohol is known to act as a stabiliser.

Thus, a typical polymerisation procedure will commence by forming an aqueous solution of the hydroxy functional polymer which acts as stabilising agent, together with a polymerisation initiator, in a reaction vessel. Then while agitating the contents of the reaction vessel, the organic monomers are progressively fed in so that the monomers become dispersed in the aqueous phase and polymerise therein. The addition of monomers can be continued until the mixture in the reaction vessel is a slurry of polymer beads containing about 30% by weight of polymer.

In a possible variation of this procedure some of
the monomer is dispersed in the aqueous solution of
stabilising agent before any polymerisation initiator is
added. In another possible variation the monomers are
emulsified in water before they are added to the reaction
vessel.

- Suspension polymerisation typically produces polymer beads with a diameter larger than  $100\mu$ . Smaller particle sizes in the range of  $50\text{--}100\mu$  can be obtained by increasing the amount of stabiliser in the aqueous phase, or by increasing the amount of agitation, or both.
- 25 Polymerisation may be carried out using a combination of polyvinyl alcohol and a second stabilising agent which may or may not be a second hydroxy-functional polymer. Examples of materials which can serve as a

13

second stabilising agent include sodium oleate and sodium lauryl sulphate, both of which are anionic surfactants, also nonionic surfactants with HLB of 8 and above, such as  $C_{13}/C_{15}$  alcohol ethoxylated with 3 to 11 ethylene oxide residues, and sorbitan ester surfactants of similar HLB.

The amount of hydroxy functional polymer present in solution as a stabilising agent will generally be between 1% and 5% by weight of the aqueous phase in which polymerisation occurs. Larger amounts can also be used.

The amount may well be between 3% and 15% by weight of the monomers undergoing polymerisation, possibly between 5% and 10%.

Observation by microscopy of such particles showed the hydroxy functional polymer to form a coating or incomplete coating of the particles, from which it can be estimated that the amount of hydroxy functional polymer attached to the particles was at least 1% by weight of these particles after (and also of course before) the incorporation of perfume.

Suspension polymerisation can be used to produce linear polymers, or cross linked polymers. The presence or absence of cross linking is determined by the monomers which are used.

As mentioned above, a molecule with more than one carbon-carbon double bond can serve as a cross linking agent. When used, a suitable amount of such crosslinking agent is not over 5 mole% of the monomer mixture, e.g. in a range from 0.5 to 3 mole%.

Hydroxy substituted acrylate esters can also lead to cross-linking. The mechanism by which they do so is a side reaction which is not fully understood. When used, a suitable amount may lie in a range from 3 to 30 mole% of the monomer mixture. Preferably 10 to 30 mole%.

After the manufacture of the particles by polymerisation, the direct product is in the form of an aqueous slurry. If desired, the particles may be separated from the aqueous phase by filtration or centrifuging, possibly followed by drying.

Another possible route for the production of polymer particles is emulsion polymerisation to yield an aqueous emulsion of very small polymer particles (typically of sub-micron size) followed by a drying step to agglomerate these particles into larger particles with a size of at least  $20\mu$ .

Absorption of perfume by the particles can be brought about simply by bringing the perfume and the particles into contact, and allowing them to stand. This may be done by mixing perfume with the particles after they have been separated from the aqueous phase, or it may be done by mixing perfume into an aqueous slurry of the particles and allowing the mixture to equilibrate. It can be done by mixing the particles and perfume separately into an aqueous liquid product and allowing that mixture to equilibrate.

# Encapsulated particles

10

15

20

25

15

A further possibility is to encapsulate a "core" of polymer as described above, with aminoplast resin, while providing hydroxy-functional polymer at the exterior of the capsules, and absorb perfume within the core.

5 Several typical procedures are available to produce such encapsulated polymer. One procedure is to form polymer beads, for example of an acrylate polymer, as described earlier, and dispense this organic mixture in an aqueous solution containing the hydroxy functional polymer and urea-formaldehyde precondensate. The mixture 10 is agitated to keep the organic mixture in suspension. While maintaining solution temperature between 20°C and 90°C, acid is then added to catalyse polymerisation of the dissolved urea-formaldehyde precondensate. If the solution is rapidly agitated during this polymerisation 15 step, shells of water-insoluble, urea-formaldehyde polymer form around and encapsulate the dispersed organic mixture and molecules of the hydroxy-group containing polymer are incorporated in and at the exterior of these 20 shells. Melanine-formaldehyde precondensate can be used in place of urea-formaldehyde, and may be preferred.

Another procedure is to form encapsulated core polymer, in the absence of perfume, and subsequently allow perfume to diffuse through the shell, into the core polymer. We have found that absorption of perfume is possible through a thin shell, even though a thicker hollow shell is capable of retaining liquid perfume. Suitably the weight of polymer forming the shell is less

25

than the weight of polymer forming the core, and the shell to core weight ratio may lie in a range from 1:3 to 1:20, better 1:5 to 1:20.

Hydroxy functional polymer will generally provide a substantial proportion of the shell, yet constitute from 1% to 25% of the capsules.

For this procedure it has been found preferable to encapsulate monomer within an aminoplast shell, then polymerise the monomer to form a (preferably solid) core of polymer within the shell. Less preferred is to partially polymerise the core before encapsulation.

It is also possible to encapsulate a mixture of liquid monomer and fragrance, then polymerise. However, this necessarily exposes perfume to the polymerisation reaction, whereas absorption of perfume through the shell into a previously polymerised core does not.

When a product contains particles in which perfume is absorbed within polymer which is encapsulated by a thin shell, perfume can diffuse through the shell, and can be released without rupture of the shell, although the release and dispersion of perfume will be slower than for neat perfume. Thus, encapsulated polymer with absorbed perfume can provide deposition and retarded release of perfume similarly to the (preferred) arrangement when perfume is absorbed in polymer beads which have hydroxy functional polymer directly at their exterior.

17

#### Perfume

10

As is well known, a perfume normally consists of a mixture of a number of perfumery materials, each of which has a fragrance. The number of perfumery materials in a 5 perfume is typically ten or more. The range of fragrant materials used in perfumery is very wide; the materials come from a variety of chemical classes, but in general are water-insoluble oils. In many instances, the molecular weight of a perfumery material is in excess of 150, but does not exceed 300.

The perfumes used in the present invention can be mixtures of conventional perfumery materials. Perfumery materials which may be used include: acetyl cedrene, 4acetoxy-3-pentyltetrahydropyran, 4-acetyl-6-t-butyl-1,1-15 dimethylindane, available under the trademark "CELESTOLIDE", 5-acetyl-1,1,2,3,3,6-hexamethylindane, available under the trademark "PHANTOLIDE", 6-acetyl-1isopropyl-2,3,3,5-tetramethylindane, available under the trademark "TRASEOLIDE", alpha-n-amylcinnamic aldehyde, amyl salicylate, aubepine, aubepine nitrile, aurantion, 2-t-butylcyclohexyl acetate, 2-t-butylcyclohexanol, 3-(pt-butylphenyl)propanal, 4-t-butylcyclohexyl acetate, 4-tbutyl-3,5-dinitro-2,6-dimethyl acetophenone, 4-tbutylcyclohexanol, benzoin siam resinoids, benzyl benzoate, benzyl acetate, benzyl propionate, benzyl salicylate, benzyl isoamyl ether, benzyl alcohol, bergamot oil, bornyl acetate, butyl salicylate, carvacrol, cedar atlas oil, cedryl methyl ether, cedryl

acetate, cinnamic alcohol, cinnamyl propionate, cis-3-hexenol, cis-3-hexenyl salicylate, citronella oil, citronellol, citronellonitrile, citronellyl acetate, citronellyloxyacetaldehyde, cloveleaf oil, coumarin, 9-

- decen-1-ol, n-decanal, n-dodecanal, decanol,
  decyl acetate, diethyl phthalate, dihydromyrcenol,
  dihydromyrcenyl formate, dihydromyrcenyl acetate,
  dihydroterpinyl acetate, dimethylbenzyl carbinyl acetate,
  dimethylbenzylcarbinol, dimethylheptanol,
- dimethyloctanol, dimyrcetol, diphenyl oxide,
  ethyl naphthyl ether, ethyl vanillin,
  ethylene brassylate, eugenol, geraniol, geranium oil,
  geranonitrile, geranyl nitrile, geranyl acetate,
  1,1,2,4,4,7-hexamethyl-6-acetyl-1,2,3,4-
- 15 tetrahydronaphthalene, available under the trademark
  "TONALID",
  - 1,3,4,6,7,8-hexahydro-4,6,6,7,8,8-hexamethylcyclopenta-2-benzopyran, available under the trademark "GALAXOLIDE", 2-n-heptylcyclopentanone,
- 3a,4,5,6,7,7a-hexahydro-4,7-methano-1(3)H-inden-6ylpropionate, available under the trademark
  "FLOROCYCLENE",
  - 3a,4,5,6,7,7a-hexahydro-4,7-methano-1(3)H-inden-6-ylacetate, available under the trademark "JASMACYCLENE",
- 4-(4'-hydroxy-4'-methylpentyl)-3-cyclohexenecarbaldehyde, alpha-hexylcinammic aldehyde, heliotropin, Hercolyn D, hexyl aldone, hexyl cinnamic aldehyde, hexyl salicylate, hydroxycitronellal, i-nonyl formate,

- 3-isocamphylcyclohexanol, 4-isopropylcyclohexanol,
  4-isopropylcyclohexyl methanol, indole, ionones, irones,
  isoamyl salicylate, isoborneol, isobornyl acetate,
  isobutyl salicylate, isobutylbenzoate, isobutylphenyl
  acetate, isoeugenol, isolongifolanone, isomethyl ionones,
  isononanol, isononyl acetate, isopulegol, lavandin oil,
  - lemongrass oil, linalool, linalyl acetate, LRG 201,

    1-menthol, 2-methyl-3-(p-isopropylphenyl)propanal,

    2-methyl-3-(p-t-butylphenyl)propanal, 3-methyl-2-pentyl-
- 10 cyclopentanone, 3-methyl-5-phenyl-pentanol, alpha and beta methyl naphthyl ketones, methyl ionones, methyl dihydrojasmonate, methyl naphthyl ether, methyl 4-propyl phenyl ether, Mousse de chene Yugo, Musk ambrette, myrtenol, neroli oil, nonanediol-1,3-diacetate, nonanol,
- nonanolide-1,4, nopol acetate, 1,2,3,4,5,6,7,8-octahydro-2,3,8,8-tetramethyl-2-acetyl-naphthalene, available under the trademark "ISO-E-SUPER", octanol, Oppoponax resinoid, orange oil, p-t-amylcyclohexanone,
  - p-t-butylmethylhydrocinnamic aldehyde, 2-phenylethanol,
- 20 2-phenylethyl acetate, 2-phenylpropanol, 3phenylpropanol, para-menthan-7-ol, para-t-butylphenyl methyl ether, patchouli oil, pelargene, petitgrain oil, phenoxyethyl isobutyrate, phenylacetaldehyde diethyl acetal, phenylacetaldehyde dimethyl acetal, phenylethyl
- n-butyl ether, phenylethyl isoamyl ether,

  phenylethylphenyl acetate, pimento leaf oil,

  rose-d-oxide, Sandalone, styrallyl acetate, 1,1,4,4
  tetramethyl-6-acetyl-7-ethyl-1,2,3,4-

tetrahydronaphthalene, available under the trademark

"VERSALIDE", 3,3,5-trimethyl hexyl acetate, 3,5,5
trimethylcyclohexanol, terpineol, terpinyl acetate,

tetrahydrogeraniol, tetrahydrolinalool, tetrahydromuguol,

tetrahydromyrcenol, thyme oil,

trichloromethylphenylcarbinyl acetate, tricyclodecenyl

acetate, tricyclodecenyl propionate, 10-undecen-1-al,

gamma undecalactone, 10-undecen-1-ol, undecanol,

vanillin, vetiverol, vetiveryl acetate, vetyvert oil,

acetate and propionate esters of alcohols in the list

above, aromatic nitromusk fragrances, indane musk

fragrances, isochroman musk fragrances, macrocyclic

ketones, macrolactone musk fragrances, and tetralin musk

fragrances.

Perfumes frequently include solvents or diluents, for example: ethanol, isopropanol, diethylene glycol monoethyl ether, dipropylene glycol, diethyl phthalate and triethyl citrate.

Perfumes which are used in this invention may, if

20 desired, have deodorant properties as disclosed in US-A4303679, US-A-4663068 and EP-A-545556.

If the polymer particles are solid particles which are impregnated with liquid after manufacture, we have found that the absorption of liquid can be enhanced by choosing materials with a hydrophobic character or mixing a hydrophobic oil into the perfume.

Examples of hydrophobic oils which can enhance perfume uptake are dibutylphthalate, alkane mixtures such as

25

21

isoparaffin and  $di(C_8-C_{10} \text{ alkyl})$  propylene glycol diester.

#### Liquid properties

When the preferred solid particles are allowed to absorb liquid, they can absorb a surprising quantity, often in excess of their own weight.

We prefer to use a polymer : liquid weight ratio in a range from 4:1 to 1:10, especially from 2:1 or 3:2 up to 1:3 or 1:4.

When perfume is encapsulated in microcapsules, for instance capsules of aminoplast resin, the weight ratio of shell polymer to perfume may lie in a range from 1:30 or 1:20 to 1:2 or 1:1.

#### 15 Perfume release

If the polymer particles are solid particles impregnated with perfume, then after deposition of the particles onto a surface, such as fabric surface, hair, skin, glass etc, the perfume will be released from the surface by evaporation from the particles. We have observed that the profile of evaporation from the particles is similar to the profile of evaporation from neat perfume oil itself. As is usual, the more volatile "top note" materials evaporate first. In consequence the character of the fragrance is not greatly changed by absorption into the polymer particles.

However, we have observed that some perfume components are retained in the deposited particles during

22

the drying, whereas they are lost by evaporation during drying if deposited directly on fabric without polymer present. Thus the polymer particles may alter the character of the fragrance retained on a surface after drying, so that it more closely resembles the original perfume oil.

If the polymer particles are capsules containing liquid perfume, the perfume may be trapped within the capsules until the capsules rupture (although we prefer capsules with a thin wall, to allow the perfume to diffuse out through the intact wall). Rupture of capsules may take place during drying, notably during tumble drying of fabrics, or it may take place during handling and ironing of fabrics.

15

20

### Fabric detergent composition

Perfume-carrying particles of this invention may be incorporated into a detergent composition for fabric washing. Such a composition may be in solid form, notably a particulate or compressed solid composition, or may be in liquid form, notably with an aqueous or non-aqueous liquid phase with or without suspended solid.

Surfactants useful as detergent active in the detergent compositions herein include well-known anionic, nonionic, amphoteric and zwitterionic surfactants.

Typical of these are the alkyl benzene sulphonates, alkyl sulphonates, alkyl- and alkyl ether sulphates, primary alkyl sulphates, alkoxylated alcohols, alpha-sulphonates

23

of fatty acids and of fatty acid esters, alkyl betaines, and alkyl polyglycosides all known in the detergent art.

Detergent active is preferably present in a quantity of at least 5% or 10% by weight of a composition, and may well be in a quantity not exceeding 50% or 40% by weight. Concentrated solid detergent compositions will generally contain detergent active in a quantity from 10% to 50% by weight of the composition. Liquid compositions will typically contain from 5% to 30% by weight surfactants.

10 Detergency builders are materials which function to soften hard water by solubilisation or other removal of calcium and to a lesser extent magnesium salts responsible for water hardness compounds, especially exemplified by sodium tripolyphosphate. A further water soluble inorganic builder compound is sodium carbonate 15 which is generally used in conjunction with a seed crystal to accelerate the precipitation of calcium carbonate. Common insoluble inorganic detergency builders are zeolites. Organic detergency builders such as sodium citrate and polyacrylate can also be used. 20 detergency builder component of a detergent composition will as mentioned generally comprise from 5 to 80%, preferably from 5 to 10% to 60% by weight of the detergent composition. As is well known, many detergent compositions avoid phosphate builders. 25

A liquid composition will typically contain 5% to 40% by weight of water-soluble builder salt, partially dissolved and partially suspended in an aqueous liquid

24

phase.

Other ingredients which are customarily included in a detergent composition, although not necessarily all together, include bleaches, bleach activators, alkaline silicate, soil release agents, anti-redeposition agents such as sodium carboxymethyl cellulose, enzymes, fabric softening agents including softening clays, fluorescent brighteners, antifoam agents or conversely foam boosters and filler such as sodium sulphate.

10 A detergent composition will generally contain from 0.1 to 3% by weight of perfume, more usually not over 1.5% or 1%, at least partially carried by perfume particles according to this invention. Perfume carrying particles of this invention are preferably added to a detergent composition after any manufacturing steps involving heat (e.g. spray drying). Thus perfume carrying particles according to this invention may be added to a detergent composition, typically by mixing them into the preformed particulate or liquid detergent composition. They may be added as dry particles, or as an aqueous slurry. If the composition is a particulate solid, the water content of a slurry may be absorbed by other constituents of the detergent composition.

#### 25 <u>Fabric conditioners</u>

Perfume-carrying particles of this invention may be incorporated in fabric conditioning products used during rinsing of fabrics. The main benefits delivered by such

25

products are softness, fragrance and anti-static. Softness is usually the most important.

A fabric softening product contains at least one softening agent which functions to give the fabric a softer handle. Frequently such agents also provide an anti-static benefit. Such agents are usually cationic, but may be nonionic, amphoteric or zwitterionic materials.

Many fabric softening products take the form of

compositions intended to be added to rinse water. The
fabric softening agents are then materials with low
solubility in water, and which deposit on the fabrics.

Typically the solubility in acidified water at 20°C is
less than 10g/litre, preferably less than 1g/litre. When

added to rinse water such materials form a dispersed
phase which is then able to deposit on fabrics which are
being rinsed in the water.

Many commercially important fabric softening agents are organic compounds containing quaternary nitrogen and 20 at least one carbon chain of 6 to 30 carbon atoms, e.g. in an alkyl, alkenyl or aryl substituted alkyl or alkenyl group with at least six aliphatic carbon atoms.

Other fabric softening agents are the corresponding tertiary amines and imidazolines, other aliphatic

25 alcohols, esters, amines or carboxylic acids incorporating a C8 to C30 alky, alkenyl or acyl group, including esters of sorbitan and esters of polyhydric alcohols, mineral oils, polyols such as polyethylene

glycol, and also clays.

Some specific instances of fabric softening agents are:

1) Acrylic quaternary ammonium compounds of the formula

5 (I)
$$Q_{1} \longrightarrow \begin{array}{c} Q_{1} \\ \downarrow \\ Q_{2} \end{array} \qquad X^{-} \qquad (I)$$

10

15

20

wherein each  $Q_1$  is a hydrocarbyl group containing from 15 to 22 carbon atoms,  $Q_2$  is a saturated alkyl or hydroxy alkyl group containing from 1 to 4 carbon atoms,  $Q_3$  may be as defined for  $Q_1$  or  $Q_2$  or may be a phenyl and X-as an anion preferably selected from halide, methyl sulphate and ethyl sulphate radicals.

Throughout this discussion of fabric softening agents the expression hydrocarbyl group refers to alkyl or alkenyl groups optionally substituted or interrupted by functional groups such as -OH, -O-, -COHN-, -COO-, etc.

Representative examples of these quaternary softeners include ditallow dimethyl ammonium chloride; ditallow dimethyl ammonium methyl sulphate; dihexadecyl dimethyl ammonium chloride; di(hydrogenated tallow) dimethyl ammonium methyl sulphate or chloride; di(coconut)dimethyl ammonium chloride dihexadecyl diethyl ammonium chloride; dibenhenyl dimethyl ammonium chloride.

Examples of commercially available materials in this

30 class are ARQUAD 2C, ARQUAD 2HT, ARQUAD 2T (all Ex Akzo
Chemie) and PRAPAGEN WK, PRAPAGEN WKT, DODIGEN 1828 (all

10

Hoechst).

2) Diamido Quaternary Ammonium Salts

Diamido quaternary salts of general formula (III)

5 are also known to be useful as fabric softening agents.

$$\begin{array}{c|cccc}
O & Q_{2} & O \\
\parallel & & \parallel & \parallel \\
Q_{1}-C-NH-Q_{6}-N^{+}-Q_{6}-NH-C-Q_{1} & X^{-} \\
& & \downarrow & \\
Q_{5}
\end{array} (III)$$

 $Q_6$  is a divalent alkylene group containing from 1 to 3 carbon atoms.  $Q_1,\ Q_2,\ Q_5$  and X are as defined previously.

15 Examples of suitable materials are methylbis
(tallowamidoethyl)(2-hydroxyethyl) ammonium methyl
sulphate and methyl bis (hydrogenated tallowamido
ethyl)(2 hydroxyethyl) ammonium methyl sulphate. These
materials are available from Sherex Chem Co under trade
20 names VARISOFT 222 and VARISOFT 110 respectively and
under the trade name ACCOSOFT from Stepan.

3) Ester Quaternary Ammonium Salts

A number of ester groups containing quaternary

25 ammonium salts, including those disclosed in EP 345842 A2

(Procter), EP 239910 (Procter) and US 4137180 (Lever) and incorporated herein by reference, are known to be particularly useful as softening materials. These materials can be represented by generic formulae (IV) and

30 (V) below.

28

In formula (IV) Q<sub>7</sub> is a hydrocarbyl group containing 1 to 4 carbon atoms, Q<sub>8</sub> is  $(CH_2)_n$ -Z-Q<sub>10</sub> where n is an integer from 1 to 4 or -Q<sub>10</sub>. Q<sub>9</sub> is an alkyl or hydroxyalkyl group of 1 to 4 carbon atoms, or is as defined for Q<sub>8</sub>. Q<sub>10</sub> is a hydrocarbyl group containing from 12 to 22 carbon atoms and Y can be -CH(OH)-CH<sub>2</sub>- or Q<sub>6</sub>, as previously defined. Z can be -O-C(O)O-, -C(O)O-C(O)-O or -O-C(O)- and X<sup>-</sup> is an anion.

In formula (V) the symbols  $Q_2$ ,  $Q_{10}$ , Z and  $X^-$  have 25 the meanings defined previously.

Suitable materials of formula (IV) are N,N-di (stearyl-oxyethyl)-N,N-dimethyl ammonium chloride and N,N-di (stearyl-oxyethyl)-N-hydroxyethyl-N-dimethyl ammonium chloride. Stearyl may be replaced with oleyl, palmityl or tallowyl (mixed chain length) groups. An illustrative example of a formula (V) material is 1,2-ditallowyloxy-3-trimethyl ammoniopropane chloride, which is a ditallow ester of 2,3-dihydroxy propane trimethyl ammonium chloride (HOECHST).

35

29

4) Quaternary Imidazolinium Salts

A further class of cationic softener materials is the imidazolinium salts of generic formula (VI).

5

10

Wherein  $Q_{11}$  is a hydrocarbyl group containing from 6 to 24 carbon atoms, G is -N(H)-, or -O-, or  $NQ_2$ , n is an integer between 1 and 4, and  $Q_7$  is as defined above.

Preferred imidazolinium salts include 1-methyl-1
(tallowylamido) ethyl-2tallowyl-4,5 dihydro imidazolinium methosulphate and 1-methyl-1-(palmitoylamido) ethyl-2octadecyl-4,5-dihydroimidazolinium chloride.

Representative commercially available materials are VARISOFT 475 (Sherex) and REWOQUAT W7500 (Rewo).

20

5) Zwitterionic Fabric Softeners

Other useful ingredients of softening systems include zwitterionic quaternary ammonium compounds such as those disclosed in EP 332270 A2 (Unilever)

25 incorporated herein by reference. Representative materials in this class are illustrated by general

30

formula (XI) and (XII)

$$\begin{array}{c}
Q_{19} \\
\downarrow \\
Q_{11}-N^{+}-CH_{20}-Z \\
\downarrow \\
Q_{19}
\end{array} (XI)$$

10
$$\begin{array}{c|c}
O & Q_{19} \\
\parallel & \downarrow \\
Q_{11}-C-NH-Q_{20}-N^{+}-Q_{20}-Z^{-} \\
\downarrow & \downarrow \\
Q_{19}
\end{array} (XII)$$
15

wherein the groups  $Q_{19}$  are selected independently from  $Q_7$ ,  $Q_{11}$  and  $Q_{14}$ ;  $Q_{20}$  is a divalent alkylene group containing 1 to 3 carbon atoms and may be interrupted by -O-, -COHN, -C(O)O-, etc; and  $Z^-$  is an anionic water solubilising group (e.g. carboxy, sulphate, sulpho or phosphonium).

Examples of commercially available materials are the
25 EMPIGEN CD and BS series (Albright Wilson) the REWOTERIC

AM series (Rewo) and the Tegobetain F, H, L and N series
(GOLDSCHMIDT).

# 6) Nonionic Ingredients

It is well known to blend nonionic materials with cationic, amphoteric or zwitterionic softening materials as a means of improving dispersion of the product in rinse waters and enhancing the fabric softening properties of the softener blend.

31

Suitable nonionic adjuncts include lanolin and lanolin derivatives, fatty acids containing from 10 to 18 carbon atoms, esters or fatty acids containing from 8 to 24 carbon atoms with monohydric alcohols containing from 5 1 to 3 carbon atoms, and polyhydric alcohols containing 2 to 8 carbon atoms such as sucrose, sorbitan, together with alkoxylated fatty acids, alcohols and lanolins containing an average of not more than 7 alkylene oxide groups per molecule. Suitable materials have been 10 disclosed in EP-A-88520 (Unilever), EP-A-122141 (Unilever), GB 2157728A (Unilever), GB 8410321 (Unilever), EP-A-159918 (Unilever), EP-A-159922 (Unilever) and EP-A-79746 (Procter).

Fabric softening compositions generally do not

15 contain anionic detergent active nor bleach, nor

detergency building. It is desirable that the amounts

(if any) of anionic detergent active, bleach and

detergency builder are all less than the amount of the

fabric softening agent.

A fabric softening composition which is intended to be added to rinse water may be in the form of a solid, a powder or tablet for instance, which disperses in the rinse water.

More commonly, a fabric softening composition for
25 addition to rinse water is in the form of a liquid, and
is an aqueous dispersion in water. Such a fabric

softening composition may contain from 1% or 2% up to 30% or 40% by weight of a fabric softening agent but may contain higher levels from 40% up to 80% or even 90% by weight in a very concentrated product. The composition will usually also contain water, which may provide the balance of the composition.

Liquid fabric softening compositions are customarily prepared by melting the softening ingredients and adding the melt to hot water, with agitation to disperse the water-insoluble ingredients.

Perfume-carrying particles according to this invention may be added as dry particles or as an aqueous slurry, suitable after the composition has cooled.

10

The amount of perfume incorporated in a fabric softening product will lie in the range from 0.01% to 10% by weight.

For fabric conditioning liquids containing less than 40% by weight of fabric softening agent, the amount of perfume is preferably 0.1 to 3% by weight, more preferably 0.1 to 1.5%.

Another form of fabric softening product has a fabric softening agent in a composition which is coated onto a substrate, usually a flexible sheet or sponge, which is capable of releasing the composition in a tumble dryer. Such a product can be designed for single usage or for multiple uses. One such multi-use article

33

comprises a sponge material releasably enclosing enough of the conditioning composition to effectively impart fabric softness during several drying cycles. The multiuse article can be made by filling a porous sponge with 5 the composition. In use, the composition melts and leaches out through the pores of the sponge to soften and condition fabrics. A single use sheet may comprise the inventive compositions carried on a flexible substrate such as a sheet of paper or woven or non-woven cloth 10 substrate. When such an article is placed in an automatic laundry dryer, the heat, moisture, distribution forces and tumbling action of the dryer removes the composition from the substrate and deposits it on the fabrics. Substrate materials for single use and multiple use articles, and methods of impregnating or coating them are discussed in US-A-5254269 and elsewhere.

A fabric softening product which is an impregnated or coated sheet, sponge or other substrate will typically contain perfume-carrying particles in a quantity to

20 provide from 0.5 to 8% by weight perfume, preferably from 2% or 3% up to 6%.

#### Personal washing products

Perfume-containing particles in accordance with this

25 invention can be utilised in soap bars and in so-called

syndet bars which contain non-soap detergent active, but

34

are used for personal washing. Sodium fatty acyl isethionate is commonly used in this context. In such products the particles may provide from 0.5 to 5% by weight perfume.

Perfume-containing particles in accordance with this invention can be utilised in liquid products for personal washing, such as hair shampoos or shower gels. These typically contain from 2 to 40% by weight of detergent active which is anionic, amphoteric, nonionic or a mixture of these, especially a mixture in which anionic and/or amphoteric detergent active provides 2 to 40% by weight of the composition.

Such products will frequently include other materials, especially 0.01 to 3% by weight of cationic polymer and/or 0.1 to 10% by weight of silicone to deposit on skin or hair.

15

Again in such products the particles may provide from 0.5 to 5% by weight perfume.

Preferably such liquid compositions will be of

20 desirable viscosities, so as to be pourable or
squeezable, and yet will not be so thin that they run
uncontrollably. The desired viscosity range is
approximately 1,000 to 15,000 centipoises at room
temperature (25°C) and low shear rate, preferably 3,000

25 to 6,000 centipoises at 10 sec-1 shear rate.

35

#### Household Cleaning Products

The perfume-containing particles of the present invention can also be utilised in household products for cleaning hard surfaces. These are usually aqueous

5 liquids containing detergent active and/or polymeric thickener. Other common ingredients are solvents and bleach. The amount of detergent active which is used may vary widely. For some products a low level such as 0.5-3% is appropriate, while others may have more, ranging up to 15% or 20% by weight. Product viscosity will frequently exceed 100 centipoise at 10 sec<sup>-1</sup> shear rate and a range of 200 to 2000 centipoise at 10 sec<sup>-1</sup> shear rate is common.

#### 15 <u>Sunscreening Compositions</u>

The particles may imbibe sunscreening agents, where necessary after solution in a carrier oil, such as benzophenones, methyl anthranilate, methoxycinnamates, octocrylene, phenyl salicylates, paraamino benzoyl esters, homosalate, benzylidene camphor, phenylbenzimiolazol sulphonic acid and dibenzoylmethane.

# "Leave-on" Hair Care, Skin Care, Fabric and Household Compositions

Particles with imbibed perfumes may be used in "leave-on" compositions such as hand-creams, skin-creams

and lotions, foundation creams, under-arm deodorants and anti-perspirants, in otherwise conventional aerosol, roll-on, cream and stick formulations, and leave-on hair-conditioners. They may also be formulated in such fabric and household products as fabric fresheners, carpet fresheners, ironing aids and toilet rim blocks.

The invention will be further explained and exemplified by the following Examples in which all parts and percentages are by weight, unless stated otherwise.

#### 10 Example 1

A 700ml reaction flask equipped with motorised stirrer, reflux condenser, thermometer and inlet tube for delivery from a peristaltic pump was placed in a water bath at about 65°C.

- An aqueous phase was prepared by mixing hydroxyethyl cellulose (5 parts) and deionized water (168 parts). The hydroxyethyl cellulose had a degree of substitution of one and was available from Hercules Chemical Corp as NATROSOL 250L. This phase was mixed until the
- 20 hydroxyethyl cellulose dissolved and was then charged into the reaction flask. Stirring was applied to the reaction flask.

A monomer phase was prepared by mixing iso-butyl methacrylate (70 parts) with a cross linking co-monomer which was 1,6-hexanedioldiacrylate (1.8 parts).

2,2'-anzo(bis)isobutyronitrile [usually abbreviated

37

to AIBN] (2 parts) was added directly to the reaction flask and dispersed for about five minutes.

The monomer phase was added to the stirring reaction flask using a peristaltic pump over about ninety minutes.

5 After addition the reaction mass was stirred at about 65°C for about three hours and subsequently cooled.

The polymer beads were recovered from the aqueous slurry by filtration and air dried. The beads were sieved to separate the fraction with size below 125µm.

It can be seen that in this Example, the total quantity of hydroxy-functional polymer was less than 10% by weight of monomers, and only part of that quantity becomes attached to the polymer beads.

The cross linking agent is 2% by weight and 1.63% by mole of the monomer mixture.

## Example 2

A 700ml reaction flask, equipped as in Example 1, was placed in a water bath at about 65°C.

An aqueous phase was prepared by mixing poly(vinyl alcohol) available as Gohsenol AH-22 from Nippon Gohsei British Trades and Shippers Ltd, Dagenham, Essex and having a degree of hydrolysis of 97 to 98.8% (5 parts) and deionized water (168 parts). This phase was stirred until the poly(vinyl alcohol) dissolved and was then charged into the reaction flask. Stirring was applied to

the reaction flask.

A monomer phase was prepared by mixing styrene (68 parts) and 1,6-hexanedioldiacrylate (1.8 parts).

AIBN (2 parts) was added directly to the reaction flask and dispersed for about five minutes.

The monomer phase was added to the stirring reaction flask using a peristaltic pump over about ninety minutes. After addition the reaction mass was stirred at about 65°C for about three hours and subsequently cooled.

At this stage, the products was in the form of an aqueous slurry from which the polymer beads were recovered by filtration and air dried.

#### Examples 3 to 5

The procedure of Example 2 was repeated with different monomers as follows:

Example 3: Methyl methacrylate (70 parts) and 1,6-hexanedioldiacrylate (1.8 parts)

Example 4: n-Butylmethacrylate (70 parts) and 1,6-hexanedioldiacrylate (1.8 parts)

20 Example 5: iso-butylmethacrylate (54 parts) and hydroxypropylmethacrylate (18 parts)

In each of these examples the monomers were satisfactorily converted to polymer beads. These were recovered by filtration and air dried.

#### Example 6

Beads, produced as in Example 5, were sieved to remove any beads larger than  $75\mu m$  diameter.

A perfume was prepared consisting of equal amounts of

- 5 i) dihydromyrcenol (2,6-dimethyl-7-octen-2-ol)
  - ii) anisaldehyde
  - iii) dimethylbenzylcarbinyl acetate
- (iv) 1,1,2,4,4,7-hexamethyl-6-acetyl-1,2,3,4tetrahydronaphthalene, available under the trademark
  "TONALID",
  - (v) 3a,4,5,6,7,7a-hexahydro-4,7-methano-1(3)H-inden-6ylpropionate, available under the trademark
    "FLOROCYCLENE",
- Perfume-loaded polymer beads were prepared by mixing the above beads and perfume into a diluted rinse conditioner, to yield an aqueous slurry containing:-

	Polymer beads	10.71%
20	Perfume	10.71%
	Dihardened tallow dimethyl	
	ammonium chloride	3.5%
	Water	Balance

This slurry was agitated for two hours and left to stand for twenty four hours, after which it appeared that

40

all the perfume had been absorbed into the polymer beads.

This slurry was added to a quantity of a rinse conditioner formulation which was an aqueous emulsion containing a 1,2-dihardened tallowloxy-3-trimethyl ammoniopropane chloride (HTTMAPC) as cationic softener. This material is disclosed in US-A-4137180.

The formulation contained:

	HTTMAPC (including some fatty acid impurity	13.5% y)
	Ethoxylated Coconut alcohol (20E0)	0.75%
5	Hardened tallow alcohol	0.75%
	Calcium chloride	0.2%
	Preservative	0.02%
	Demineralised water	Balance to 100%

After adding the slurry, the resulting perfumed formulation contained 0.75% by weight perfume, carried in polymer beads. The perfumed rinse conditioner formulation was agitated for two hours and then stored for six days in a closed container. A control

15 formulation contained 0.75% by weight perfume, and the same concentration of fabric softener, without polymer beads.

To test perfume deposition, this rinse conditioner formulation and the control formulation were both diluted with water to provide rinse liquors containing 0.5% of the rinse conditioner formulation.

Test pieces of fabric were de-sized cotton terry towelling, approximate weight 25g. For each test, a piece of terry towelling was weighed accurately and treated with 30 times its own weight of rinse liquor, in a Tergotometer pot, stirring at 80rpm for 20 minutes. The cloth was then wrung out by hand, and line dried.

The amount of perfume in the fresh and used rinse liquors was determined by solvent extraction from 100g of rinse liquor and gas chromatographic (GC) analysis of the solvent extract. The percentage deposition of perfume materials onto the cloth was calculated for three of the five materials. The results obtained were:

	Ingredient	nt % Deposition	
		Control	Perfume in polymer beads
10	Dihydromyrcenol	14	25
	DMBCA	24	33
	Florocyclene	38	42

The amount of perfume on the dry cloth was determined by extraction of 5g of dry cloth with 20ml ethyl acetate, followed by GC analysis of the solvent extract.

The amount of perfume detected was expressed as a 20 percentage of the theoretical maximum quantity (which would be present if there were complete deposition onto fabric and no subsequent losses).

The results obtained for materials were:

43

	% Ingredient remaining on dry cloth	
Ingredient	Control	Perfume in polymer beads
Dihydromyrcenol	not detected	1.3
DMBCA	not detected	8.0
Florocyclene	not detected	9.8
Tonalid 2	30.2*	51.1*

<sup>\* =</sup> result possibly affected by other GC peaks.

10

5

#### Example 7

Perfume mixture was incorporated into polymer beads, as set out in the previous Example, but with two ratios of polymer to perfume. These were 1:1 polymer to perfume and 1:2 polymer to perfume.

The resulting perfumed beads were incorporated into rinse conditioner formulations as in the previous example so as to provide 0.75% by weight perfume in each formulation. A control formulation contained 0.75% by weight perfume, but no polymer. These were diluted to rinse liquors containing 0.5% by weight fabric conditioner, and used to treat terry towelling as in the previous Example.

The treated cloths were assessed by a panel of eight 25 people.

Assessments were made on damp cloth directly after treatment, on dry cloth 24 hours after treatment and on

44

dry cloth five days after treatment.

The assessments were:

5	Assessment Stage	Control (no polymer)	1:1 polymer to perfume	1:2 polymer to perfume
	Damp Cloth	Strongest	Weaker than control Equal to (1:2)	Weaker than control Equal to (1:1)
	Dry Cloth (24 hours	Intense Tonalid odour	Florocyclene, Tonalid and anisaldehyde odours Stronger than (1:2)	Florocyclene, Tonalid and anisaldehyde odours Weaker than (1:1)
10	Dry Cloth (five days)	Tonalid odour	Florocyclene, Tonalid and anisaldehyde odours Weaker than (1:2)	Florocyclene, Tonalid and anisaldehyde odours Stronger than (1:1)

#### Example 8

Beads were produced as in Example 2. The monomer

15 mixture was isobutyl methacrylate (70 parts) mixed with

1,6-hexanedioldiacrylate (1.8 parts). Preparations were
carried out using various different grades of polyvinyl
alcohol and some other materials as the stabiliser. The
grades of polyvinyl alcohol differed in the extent to

20 which they had been hydrolysed from polyvinyl acetate.

45

The	materials	used	were:
-----	-----------	------	-------

		Stabiliser	Hydrolysis	Tradename
	A	polyvinyl alcohol	98-99%	Gohsenol N-300
	В	polyvinyl alcohol	97-98.5%	Gohsenol A-300
5	С	polyvinyl alcohol	87-89%	Gohsenol GH-23
	D	polyvinyl alcohol	78-80%	Gohsenol KH-17
	E	polyvinyl alcohol jointly with a surface active acrylamide copolymer	97-99%	Gohsenol AH-22
	Ĥ	polyvinyl alcohol jointly with a 95:5 copolymer of acrylamide and behenyl (25 ethylene oxide) methacrylate	97-99%	Gohsenol AH-22
	G	hydroxyethyl-cellulose		Natrosol 250L
10	Н	surface active acrylamide copolymer (comparative example)		MER 10

In all cases the dried beads were mixed with their own weight of a single perfumery material, Florocyclene, also used in Example 6. After mixing they were left to equilibrate overnight, then a small quantity of finely divided silica (2% based on the total weight of perfume and polymer) was added as a surface improving flow aid.

The perfume-containing polymer beads were mixed into 20 rinse conditioner (as used in Example 6) using a magnetic stirrer. In each case the amount of perfume carrying beads was 1.5% by weight of the rinse conditioner, so that the quantity of perfume was 0.75% by weight of the

46

rinse conditioner formulation. A control formulation was prepared containing 0.75% Florocyclene without the polymer.

Deposition onto fabric was assessed using test

5 cloths made of cotton terry towelling, acrylic fabric,
and polyester. The fabric washing procedure was the same
as set out in Example 6. After treatment of the fabric
and drying, the intensity of Florocyclene on groups of
the dried test cloths was assessed by a panel of five

10 assessors who ranked the cloths in order of intensity of
Florocyclene. The following results were obtained as
unanimous views of the assessors:

On cotton test pieces B>A>C>Control

15 C>D>Control

E>F>Control

and G>H>Control

On acrylic test pieces A>B>C>Control

On polyester test pieces A,B and C were all stronger than the control, but the assessors were not unanimous as to which of the three gave the highest perfume intensity.

These results demonstrate that the polymer particles
A,B,C and D made using polyvinyl alcohol as stabiliser
and hydroxy-functional monomer all gave an increase in
the perceived intensity of perfume on fabric. So did

47

polymer particles E and F made using polyvinyl alcohol in a mixture of stabilisers. The polymer particles G made using hydroxyethylcellulose as stabiliser and hydroxyfunctional monomer also gave an increase in the perceived intensity of perfume on fabric and were superior to the particles H made using a different polymer as stabiliser.

#### Example 9

Polymer beads were prepared by the procedure of

10 Example 2, using isobutyl methacrylate as the sole

monomer, without cross-linking agent.

Polymer beads (0.6% by weight) and dimethyl dibenzyl carbinyl acetate (DMBCA) (0.6% by weight) were added to a rinse conditioner formulation (as in Example 6). A control formulation contained 0.6% by weight of DMBCA, but no polymer beads.

After mixing the formulations, they were left in sealed containers for one week to equilibrate.

To test perfume deposition, the rinse conditioner

20 formulation and the control formulation were both diluted with water to form rinse liquors containing 0.3% by weight of the formulation.

These liquors were used to wash terry towelling test pieces, as in Example 6. After drying, the test pieces

25 were assessed for intensity of DMBCA by a panel of seven assessors. They unanimously considered that the DMBCA

was present on the cloth with more intensity when polymer beads were used, compared with the control where no polymer beads were used.

The same result was obtained using phenylethyl isoamyl ether, available under the trade mark ANTHER, in place of DMBCA.

The same result was also obtained using FLOROCYCLENE, as referred to in Example 6, in place of DMBCA.

In a similar procedure the rinse conditioner was dosed with 0.6% polymer beads and 0.2% of FLOROCYCLENE.

The control formulation was dosed with 0.6% FLOROCYCLENE only. When the dried test pieces were assessed, the perfume intensity on the cloth from 0.2% FLOROCYCLENE

15 plus 0.6% polymer beads was greater than the intensity from 0.6% FLOROCYCLENE without polymer beads.

Stability of a preferred rinse conditioner was also tested.

A perfume contained equal weights of

20 hexanal

d-limonene

phenylethyl alcohol

anisic aldehyde

dimethyl benzylcarbinyl acetate

25 2,6,6-trimethyl cyclohexadiene carboxylate (ethyl safranate)

49

5-ethoxycarbonyl-6-isopropyl bicyclo [2,2,1] hept-2-ene (herbanate)

alkyl cyclohexyl propionate

2,2-dimethyl-3(p-ethylphenyl)propanol (floralozone)

5

This perfume was mixed with and absorbed into polymer beads as above, also into beads as in Example 8G.

The weights of polymer and perfume were equal.

The perfume-loaded beads were mixed into a rinse conditioner formulation as used in Example 6, so as to incorporate 0.75% by weight perfume. As a control, the rinse conditioner formulation was mixed with 0.75% by weight of the perfume.

Samples were stored at room temperature of 20°C or at 37°C for four weeks. Then the perfume content was analysed and the condition of the products was examined. The results were

20

	Perfume rem	Perfume remaining (wt%)		earance
	20°C	37°C	20°C	37°C
Control	55.7	50.0	ok	gelled
Example 8G	64.7	52.3	ok	viscosity
				increased
Example 9	66.4	55.3	ok	viscosity
	ļ			increased

## Example 10

A laundry detergent powder base was prepared by spray drying some of its ingredients and then adding other to give the formulation tabulated below in which percentages are based on the complete formulation.

	Spray dried	<u>8w/w</u>
	Linear alkyl benzene sulphonate	7.00
	Sodium tripolyphosphate	25.00
10	Acrylic/maleic copolymer	1.00
	Sodium silicate	5.00
	Sodium sulphate	25.00
	Optical Brightener	0.20
	Chelant .	0.50
15		
	<u>Post-dosed</u>	
	Nonionic surfactant (C14-C15 E7)	4.00
	Protease enzyme	1.00
	Lipase enzyme	0.30
20	Tetraacetyl ethylene diamine (TAED)	4.00
	Sodium perborate monohydrate	16.00
	Sodium carbonate	10.00
	Suds suppressor	1.00

25 A perfume mixture contained equal amounts of: 2-methyl-3(para-t-butylphenyl)propionaldehyde which is

51

available under the trademark "LILIAL", anisic aldehyde, d-limonene, dodecyl nitrile and hexyl salicylate.

This perfume mixture was mixed with an equal weight of polymer beads prepared as in Example 5. A small quantity of finely divided silica flow aid was added so as to form a free-flowing powder. This powder was then added to the laundry detergent powder base in such a quantity that the resulting composition contained 0.4% of the perfume mixture and 0.4% of the polymer beads. A control detergent powder consisted of 0.4% of this perfume mixture added directly to the above detergent powder base.

The control powder and the test powder were used to wash samples of desized cotton terry towelling by the following procedure:

The laundry powder was dissolved at a concentration of 0.7% by weight in water at 45°C in a Tergotometer, and mixed for two minutes. Weighed samples of fabric at a wash solution to cloth ratio of 30:1 were next added to the wash liquor and washing was carried out for 20 minutes with a stirrer speed of 100 rpm. The cloths were given a single cold rinse for ten minutes at the same solution:cloth ratio, then wrung out by hand and line dried overnight.

Residual perfume was extracted from the fabric using ethyl acetate containing fluoronapthalene as an internal

standard. The amounts of two materials were determined by gas chromatography into a mass spectrometer (GC-MS) and the results obtained were:

5	Perfume Ingredient	Control (ng/µl)	Test (ng/µl)
	lilial	0.44	5.22
	hexyl salicylate	2.83	5.45

As can be seen, the use of polymer beads greatly

10 enhanced the amount of these two perfume ingredients
deposited on the fabric.

## Example 11

A liquid laundry detergent base was prepared with the following formulation:

ingredient	% by weight
sodium dodecyl benzene sulphonate	11.3
sodium lauryl ether sulphate	3.2
sodium tripolyphosphate	11.0
sodium pyrophosphate	4.0
sodium carbonate	1.75
fluorescer	0.2
silicone oil antifoam	0.5
water	balance

25

20

Polymer beads were prepared as in Examples 2 and 8. The monomer mixture was isobutyl methacrylate (70 parts) mixed with 1,6-hexanediol diacrylate (1.8 parts) as in

53

WO 98/28398 PCT/GB97/03530

Example 8. Preparations were carried out using two different grades of polyvinyl alcohol as the stabiliser. The stabilisers were:

A) Polyvinyl alcohol of 87-89% hydrolysis; Gohsenol GH-23.

5

B) Polyvinyl alcohol of 95-97% hydrolysis; Gohsenol C-500.

Samples of these polymer beads were mixed with their

own weight of FLOROCYCLENE and then the perfume

impregnated beads were mixed into the laundry liquid so

as to incorporate 0.4% by weight perfume. A control

liquid contained the perfume added directly without

polymer beads.

- These liquid compositions were used to wash desized cotton terry towelling test cloths by the following procedure:
  - 7.5 mls laundry detergent liquid was mixed with 600 mls water at  $40\,^{\circ}\text{C}$  for two minutes in a Tergotometer. A
- 20 15g piece of terry towelling was added to the
  Tergotometer pot and agitated in the pot for thirty
  minutes. The cloths were then hand wrung and line dried.
  Further cloths were washed in the same way, but rinsed in
  600 mls cold water for two minutes before wringing and
  25 line drying.

All the cloths were assessed for FLOROCYCLENE

54

intensity by a panel of four assessors. Assessment was carried out when the cloths were still damp and again after the cloths had been left to dry completely overnight.

At each assessment each assessor graded a control cloth and two test cloths assigning a score of 1 to the most intense, 3 to the least intense and 2 to the cloth of intermediate intensity. If two cloths could not be distinguished, both were given a halfway score, for instance 1.5 if the most and medium intensity cloths were indistinguishable. The results were averaged and are set out in the following table:

	Beads A	Beads B	Control
Damp, no rinse	1.5	1.5	2.75
Dry, no rinse	2.0	1.0	3.0
Damp, with rinse	2.25	1.0	2.75
Dry, with rinse	2.37	1.0	2.62

20

15

#### Example 12

Polymer beads B from the previous Example were impregnated with their own weight of FLOROCYCLENE and then treated with a small amount of finely divided silica flow aid (4% of the weight of polymer) so as to produce a free-flowing powder product. These perfume impregnated polymer beads were then used in the preparation of a hard

55

surface cleaning composition and a corresponding control formulation as set out in the following table:

	Example	Control
Citric acid	3.0	3.0
Water	92.75	93.25
Nonionic detergent	3.0	3.0
C10 alcohol 7E0	•	
Xanthan gum	0.25	0.25
Florocyclene	0.5	0.5
Beads Ex 11B	0.5	

The compositions were tested by brushing 0.46 mls composition over a white ceramic tile 15cm square. One set of tiles was left unrinsed, assessed after 1½ hours and again after leaving overnight. A second set of tiles was rinsed with 100 mils distilled water, then assessed.

Assessment was carried out by a panel of five assessors who observed that the composition with polymer 20 beads led to a more intense odour in each comparison.

#### Example 13

Polymer beads as used in Example 8G were mixed with their own weight of phenylethyl isoamyl ether (ANTHER)

25 and then treated with finely divided silica flow aid (4% of the weight of the polymer). The resulting perfume-impregnated beads were mixed with a quaternary

imidazoline fabric softener (REWOQUAT W7500/H from Sherex) so that the resulting test composition contained 4% ANTHER.

A control composition was the same quaternary softener mixed with 4% ANTHER, without polymer beads.

Test and control sheet conditioners were made by coating the test and control compositions onto non-woven fabric sheets.

Terry towelling test cloths, 20cm x 20cm, and other

laundry (to add bulk) were washed in unperfumed washing

powder and then tumble dried. Each tumble dryer

contained a test cloth; several items of other laundry,

and a sheet impregnated with either the test or the

control composition. The tumble dryers were operated on

a 50 minutes programme.

After drying, the perfume odour on the terry towelling test cloths was evaluated by a panel of five people who unaminously considered that cloths dried along with the test sheets smelt strongly of ANTHER whereas there was little or no residual ANTHER smell on the cloths dried with control sheets.

## Example 14

This example illustrates improved deposition and retention of volatile perfume on hair.

A hair shampoo was prepared with the following base

formulation:

ingredient	% by weight	
sodium lauryl ether sulphate	9.6	
ammonium lauryl ether sulphate	4.5	
sodium chloride	2.0	
citric acid	to give pH 6.0-6.5	
preservative	qs	
water	balance to 100%	

10

5

A light citrus perfume, which would not normally be substantive to hair, was prepared with the following formulation:

		<u>wt</u> %
15	Allyl amyl glycollate	3.0
	Citrathal concentrated	5.3
	Linalol	32.0
	Linalyl acetate	30.2
	Litsea Cubeba Oil	3.0
20	Orange oil	26.5

Perfume beads were prepared, generally as in Example 8G, as a 27 wt% slurry in water. This slurry was sieved to exclude any beads larger than  $125\mu m$ .

25 The citrus perfume above was added to the slurry in an amount equal to the weight of polymer in the slurry, then mixed overnight to allow the perfume to be absorbed.

58

Polymer beads according to Example 8E were also prepared as an aqueous slurry, then air dried and sieved to exclude any beads larger than 125µm. The polymer beads were mixed with finely divided silica flow aid in 5 an amount equal to 4% by weight of the beads. The above citrus perfume was then mixed with the polymer beads to produce a mixture containing equal weights of polymer and perfume. The resulting mixture was left to stand overnight in a closed container to allow absorption of the perfume.

Perfumed shampoos were prepared by mixing the above shampoo base with either of the perfumed polymer beads in sufficient quantity that the resulting shampoo contained 1% by weight perfume. A control shampoo was prepared by mixing 1% of the citrus perfume directly into the shampoo.

The shampoos were tested on hair switches by the following procedure:

Two 12g hair switches (Yugoslavian Red Tie) were used for each shampoo. Two aliquots, each of one gram, shampoo were measured onto watch glasses for each shampoo tested. The hair switch was wet under a running tap (~40°C, flow rate 40ml/sec) for 5 sec, the first aliquot of shampoo was spread along the hair and massaged into the hair for 30 sec, then the switch was rinsed for

59

20sec. The second aliquot of shampoo was then applied in a similar manner, massaging for 30sec and rinsing for 30sec. Washed switches were placed in polyethylene bags whilst other switches were washed.

5

The switches were evaluated for perfume intensity while damp, then when they were almost dry after 5 hours drying at room temperature, and again after 24 hours at room temperature. Evaluation was by a panel of four people who scored the switches 1,2 and 3 in order of perfume intensity. Their averaged scores are given in the following table:

1	5

	Control	Example 8G	Example 8E
when damp	3	2	1
after 5 hours	3	2	1
after 24 hours	2.6	2.4	1

#### 20 Example 15

Perfume was encapsulated in an aminoplast resin by the following procedure:

2.5g trimethylolmelamine condensate (Beetle resin
25 PT336 from British Industrial Plastics Ltd) was mixed
with 2.5g of polyvinyl alcohol, 2.8g sodium dihydrogen
orthophosphate and 144.7g water, to form an agueous

solution which was stirred in a beaker with a magnetic stirrer. The solution was adjusted to pH 4.3 with acetic acid, and then the solution was stirred at 20°C for 90 minutes. This aqueous phase was then mixed with 60g

5 perfume in a Silverson Mixer and homogenised to an emulsion which was stirred for 60 minutes at room temperature. The emulsion was then stirred for 20 hours at 40°C in a vessel with thermometer and condenser, then raised to 70°C for three hours. After cooling, the

10 mixture was adjusted to pH 8-10 with ammonium hydroxide. The product mixture was an aqueous slurry of aminoplast microcapsules, containing the perfume. These microcapsules incorporated the hydroxy functional polyvinyl alcohol in the aminoplast resin at the surface of the capsules, so that hydroxy groups of the polymer

The capsules were incorporated into a rinse conditioner formulation as used in Example 6, so as to provide 0.75% perfume in the rinse conditioner.

are available at the surface of the particles.

In a comparative experiment, a polyacrylamide was used in place of the polyvinyl alcohol. However, addition of the resulting capsules to the rinse conditioner caused it to coagulate.

#### 25 Example 16

Acrylate polymer encapsulated within an aminoplast

61

shell was prepared as follows:

2.5g trimethylolmelamine condensate (Beetle resin PT336 from British Industrial Plastics Ltd) was mixed 5 with 2.5g of polyvinyl alcohol, a copolymer of vinyl alcohol and vinyl amine (ratio 88:12), available from Air products, 2.8g sodium dihydrogen orthophosphate and 182.2g water, to form an aqueous solution which was stirred in a beaker with a magnetic stirrer. solution was adjusted to pH 4.3 with acetic acid, and then the solution was stirred at 20°C for 90 minutes. 25g isobutyl methacrylate, 5g hydroxypropyl methacrylate and 30g benzyl alcohol as solvent mixed together, and 0.6g of an oil-soluble initiator was added, This organic solution was mixed with the aqueous solution and homogenised to an emulsion, using a Silverson Mixer. emulsion was stirred for 60 minutes at room temperature. Next, the emulsion was stirred for 20 hours at 40°C in a vessel with thermometer and condenser, then raised to 20 70°C for three hours. After cooling, the mixture was adjusted to pH 8-10 with ammonium hydroxide. The product mixture was an aqueous slurry of aminoplast microcapsules, containing the polymer. These microcapsules incorporated the hydroxy functional 25 polyvinyl alcohol in the aminoplast resin at the surface

of the capsules, so that hydroxy groups of the polymer

62

are available at the surface of the particles.

The slurry of polymer particles, containing about 12% of acrylate core polymer was mixed with 2-methyl-3(para-t-butylphenyl) propionaldehyde (LILIAL) in an amount equal to the weight of core polymer. The mixture was stirred for 8 hours, then allowed to equilibrate for 2 days before addition to a rinse conditioner formulation as used in Example 6, so as to provide 0.75% LILIAL in the rinse conditioner.

The resulting rinse conditioner was added at a concentration of 0.3% by weight to 600ml water, and used to rinse a terry towelling test cloths, whose weight was measured accurately and close to 20g. The cloth was rinsed in a tergotometer for 20 minutes, hand wrung so that the liquid:cloth ratio was 2:1 and line dried overnight. LILIAL was extracted from the dry cloth and determined by gas chromatography.

A similar preparation and test was carried out using a copolymer with a 94:6 ratio of vinyl alcohol and vinyl 20 amine. A control test was carried out using perfume but no polymer beads.

The weights found on the cloths were:

5

Mg/g cloth				
	2 days later			
	overnight	·		
Control	1.5	Not measured		
beads with 88:12 copolymer	14.7	Not measured		
beads with 94:6 copolymer	17.7	12.6		

Particles were prepared generally as above, using

90g of an oil in place of the 30g benzyl alcohol. The

perfume odour was found to persist on the fabric during

drying, much better than a control where the perfume was

not carried by polymer particles.

#### 15 <u>Example 17</u>

A shower gel base was prepared with the following formulation:

		<u>wt</u> 8
20	Sodium lauryl ether (2EO) sulphate	16.8
	Coconut diethanolamide	1.5
	Sodium chloride	2.0
	EDTA sodium salt	0.1
	Preservative	q.s.
25	Citric acid	to pH 6 to 6.5
	Water	balance to 100%

64

A control formulation contained 1.0% by weight of a perfume, of the following composition:

		•
		<u>wt</u> %
5	Dodecanal	1.25
	Methylnonyl aldehyde	2.5
	Allyl amyl glycollate	1.5
	Anethole synthetic	0.2
	Benzyl salicylate	15.00
10	Carvone laero	0.1
	Cedramber	4.4
	cis-3-hexenyl salicylate	0.50
	Citronellol standard	2.5
	Coumarin	1.0
15	Damascone alpha	1.0
	Dihydro myrcenol	12.0
	Dipropylene glycol	6.38
•	Eugenol	0.8
	Tonalid	9.0
20	Heliotropin	0.7
	Hexyl cinnamic aldehyde	12.4
	Lavandin oil	0.7
	Lilial	8.0
	Linalol	4.0
25	Methyl dihydro jasrnonate	7.0
	Oakmoss synthetic	0.3

65

Para t-butyl cyclo hexyl acetate	6.0
Patchouli oil	2.5
Undecalactone gamma	1.2

A test formulation contained particles prepared as in Example 16, using 94:6 vinyl alcohol/vinyl amine copolymer prepared as in Example 16, containing the same perfume in a weight ratio of 1:1 perfume:core polymer. The formulation was prepared by including an aqueous slurry of the particles in a quantity of the shower gel base, so as to include 1% by weight of perfume.

The control and test shower gels were evaluated by the following procedure:

#### 15 Washing Protocol/Evaluation:

The shower gel was applied by a second person to the subject's arms. 1 gram of each shower gel was weighed onto watch glasses. The water flow and temperature from a mixer tap was set up such that the water temperature

20 was ~40°C and the flow was ~100ml/sec. The tap was left running throughout the whole experiment. The left forearm was held under the running water for ~5 seconds and the first shower gel scooped from the watch glass and applied to left forearm, keeping the arm out of the flow of water. The shower gel was lathered by rubbing it up and down the forearm for a total of 20 times, then the

66

gel was rinsed off under the flowing tap by moving the arm up and down for a total of 10 times. The skin was patted dry with a clean towel. The whole process was then repeated for the second shower gel on the right arm.

The arms were then evaluated over time for perfume odour by a panel of four expert assessors, scoring on a scale of 0 (no perfume) to 20 (very strong perfume) as follows:

	Average	Scores
Elapsed Time	Control	Test
Initial	10.0	12.7
1 hour	8.5	10.0
3½ hours	3.2	5.3
6 hours	0.3	2.2

Sunscreen example

#### 20 Example 18

For this example, polymer beads of the following composition were produced using the procedure of example 2 - isobutyl methacrylate (98 parts), 1,6-hexanedioldiacrylate (2 parts), the polyvinyl alcohol being Gohsenol GH-20. These yielded a 25wt% slurry of beads with a mean particle size of approximately 25µm into which was then stirred an equal weight of Parsol

67

MCX\* (equal to the dry weight of the polymer) to produce the particles containing sunscreen.

## \*Ex Hoffmann LaRoche

These sunscreen particles were then incorporated

into the following shampoo formulation to give a shampoo with improved deposition of sunscreen to the hair:

			wt%
	EMPICOL ESB 3	(i)	35.00
10	TEXAPON ALS	(ii)	15.00
	SODIUM CHLORIDE		1.00
	PURIFIED WATER		to 100.00
	Sunscreen/Polymerpar	cticles	1.0
	Polyquartenium-10	(iii)	0.70
15	CITRIC ACID		to pH 6.0-6.5

- (i) ex Albright & Wilson, Cumbria, England
- (ii) ex Henkel Chemicals Ltd, Henkel House, Middlesex,
  England
- 20 (iii) Celquat SC240 ex National Starch and Chemical Co.

68

#### CLAIMS:

- Particles containing organic polymer, which are insoluble in water, with a liquid absorbed in the solid
   particles, the particles having at their exterior a further polymer which incorporates free hydroxy groups.
  - 2. Particles according to claim 1 wherein the liquid is at least in part hydrophobic.

10

- 3. Particles according to claim 2 which have a core of organic polymer with liquid absorbed therein, enclosed within a polymer shell, and with said further polymer which incorporates hydroxy groups present at the exterior of the shell.
  - 4. Particles according to claim 2 or claim 3 wherein the organic polymer is a polymer of a vinyl monomer.
- 20 5. Particles according to claim 4 wherein the organic polymer is a polymer of one or more monomers which are acrylic and/or alkyl acrylic esters, of formula

69

 $H_2C = C$   $CO_2R_2$ 

5

10

where  $R_1$  is hydrogen or alkyl (including branched alkyl) of 1 to 6 carbon atoms, preferably 1 to 3 carbon atoms and  $R_2$  is alkyl (including branched alkyl) of 1 to 8 carbon atoms.

6. Particles according to any one of the preceding claims wherein said further polymer is a polymer or copolymer of vinyl alcohol.

15

- 7. Particles according to claim 6 wherein said further polymer is polyvinyl alcohol.
- Particles according to any one of claims 1 to 5
   wherein said further polymer is cellulose or chemically modified cellulose.
- Particles according to claim 5 wherein R<sub>1</sub> is hydrogen or methyl, R<sub>2</sub> is alkyl (including branched alkyl) of 3 or
   4 carbon atoms and said further polymer is polyvinyl alcohol which is at least 88% hydrolysed from polyvinyl acetate.
- 10. Particles according to any one of the preceding 30 claims with a mean size in the range 10 to  $100\,\mu m$ .

70

- 11. Particles according to any one of the preceding claims which are solid particles, wherein the liquid is a perfume.
- 5 12. A product which is a composition or article containing particles according to any one of the preceding claims, together with other ingredients.
- 13. A product according to claim 12 which is a liquid10 composition including detergent active and/or thickener.
- 14. A product according to claim 12 which is a solid laundry detergent composition for fabric washing, containing from 5 to 50% by weight of one or more detergent active compounds, and from 5 to 60% by weight of detergency builder, and from 0.1 to 1.5% by weight of perfume.
- 15. A product according to claim 12 which is a liquid
  20 laundry detergent composition for fabric washing,
  containing from 5 to 30% by weight of one or more
  detergent active compounds, from 5 to 40% by weight of
  detergency builder, water and from 0.1 to 1.5% by weight
  of perfume.

25

16. A product according to claim 12 which is a rinse

71

conditioning liquid containing from 1 to 40% by weight of a fabric softening agent as a suspension in water, and containing 0.1 to 3% by weight perfume.

- 5 17. A product according to claim 12 which is a solid substrate impregnated or coated with a fabric softening composition, and containing from 2 to 8% perfume by weight based on the fabric softening composition.
- 10 18. A product according to claim 12 which is in the form of a bar for personal washing, wherein perfume containing particles provide from 0.5 to 5% by weight of perfume.
- 19. A product according to claim 12 which is in the form of a liquid for personal washing, containing 2 to 40% by weight detergent and from 0.5 to 5% by weight of perfume.
  - 20. A product according to claim 12 which is a stay-on cosmetic product.

20

21. A sunscreen product which is a composition or article containing particles according to any one of claims 1 to 10 wherein the liquid is a sunscreening agent.

25

22. A method of making particles according to any one of

72

claims 1 to 11 which comprises, after completion of the polymerization of the particles, placing them in contact with the liquid and causing or allowing absorption of the liquid into the solid particles.

## INTERNATIONAL SEARCH REPORT

Interr mail Application No PCT/GB 97/03530

A CLASS					101/00 3/	77 03330
IPC 6	C11D17/00	A61K7/06	A61K7/48	C11D3/00	)	
ł	to International Patent Clas	sification(IPC) or to bot	th national classification r	and IPC		
	SEARCHED	and a second and the				
IPC 6	ocumentation searched (cla C11D A61K	assification system rollo	owed by classification syn	nbols)		
Documenta	tion searched other than m	inimum documentation	to the extent that such de	ocuments are includ	ed in the fields se	earched
Electronic d	data base consulted during	the international search	h (name of data base and	d, where practical, s	earch terms used	ŋ
2.20100						
	ENTS CONSIDERED TO E		<del></del>		<del></del>	
Category *	Citation of document, wit	h indication, where app	propriate, of the relevant p	passages		Relevant to claim No.
X	14 August 1	.2 A (NIPPON .991 1-8; example	CATALYTIC CHE es 1,2	EM IND)		1-7,9,11
A	WO 93 22417 A (UNILEVER PLC ;UNILEVER NV (NL)) 11 November 1993				1-13,15	
						·
<u> </u>	er documents are listed in t		кс. Х	Patent family mer	mbers are listed in	n annex.
"A" documer	egories of cited documents	e of the art which is not	, 0	ater document publish or priority date and ne	not in conflict with t	the application but
conside	ered to be of particular relevious	vance	c ir al "X" do	cited to understand th invention ocument of particular	The principle or the	eory underlying the
"L" documer which is citation	nt which may throw doubts a s cited to establish the publ or other special reason (as	lication date of another sepecified)	ir "Y" do	cannot be considered involve an inventive s ocument of particular	d novel or cannot step when the doc r relevance: the cl	be considered to cument is taken alone laimed invention
other m	nt published prior to the inte	emational filing date but	" . g	cannot be considered document is combine ments, such combina in the art.	ed with one or mor	ventive step when the pre other such docu- us to a person skilled
later tha	an the priority date claimed ictual completion of theinter		"&" do	ocument member of t		
	March 1998	Hauorai searar		ate of mailing of the i		ch report
Name and ma	ailing address of the ISA	<del></del> -	-   A	uthorized officer		
	European Patent Office NL - 2280 HV Rijswijk	e. P.B. 5818 Patentlaan	12	Minnisad onico.		
	Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016 Loiselet-Taisne, S					

1

## INTERNATIONAL SEARCH REPORT

ormation on patent family members

Interr mail Application No PCT/GB 97/03530

		, , , , ,	1017 45 377 03330	
Patent document cited in search report	Publication date	Patent family member(s)	Publication date	
EP 0441512 A	14-08-91	JP 3221582 A JP 1915308 C JP 3272766 A JP 6040894 B JP 2532711 B JP 3287505 A JP 4015286 A JP 4041583 A JP 4041584 A JP 4082803 A JP 4090814 A JP 4089883 A JP 4145904 A AU 651567 B AU 7003991 A CA 2035216 A DE 69101851 D DE 69101851 T US 5374600 A US 5641847 A JP 4214498 A	30-09-91 23-03-95 04-12-91 01-06-94 11-09-96 18-12-91 20-01-92 12-02-92 12-02-92 16-03-92 24-03-92 24-03-92 28-07-94 15-07-93 30-07-91 09-06-94 11-08-94 24-06-97 05-08-92	
WO 9322417 A	11-11-93	US 5281357 A AU 4261393 A DE 69303293 D DE 69303293 T EP 0672102 A ES 2091001 T JP 7506137 T US 5281355 A US 5385959 A ZA 9303026 A	25-01-94 29-11-93 25-07-96 21-11-96 20-09-95 16-10-96 06-07-95 25-01-94 31-01-95 29-10-94	